

IMPROVEMENTS RELATING TO ASSAY DEVICESFIELD OF THE INVENTION

5 The invention relates to assay devices for immunoassays and the like.

DESCRIPTION OF THE PRIOR ART

10 Recently, in order to increase the throughput of immunoassays, assay devices in the form of chips have been developed on which is deposited an array of localised reactive sites containing potentially many different reactive species, for example different antibodies. These reactive species react with a respective different analyte
15 in a sample supplied to the chip. Following removal of the unbonded sample, the chip can then be examined to determine the presence or absence of the respective analytes. An example of a method for preparing the reaction sites is described in more detail in GB-A-2324866 and a method for
20 analysing the substrate is described in more detail in EP-A-0902394. Both applications are incorporated herein by reference. The analysis typically involves viewing and measuring chemiluminescent radiation at the reaction sites using a low light level CCD camera system or the like.

25 A problem with these substrates is that they are small having typical dimensions 10mm x 10mm x 1mm thus making them difficult to handle. This problem is enhanced by the fact that the chips carry many small reaction sites which will be damaged if incorrectly handled. A typical chip
30 will have 100 or more such reaction sites.

 EP-A-197729, EP-A-745851 and GB-A-2147698 all disclose immunassay devices with inserts, which carry reactive sites, for location in respective wells. These are relatively cumbersome and much less convenient than chips.

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SUMMARY OF THE INVENTION

In accordance with the present invention, an assay assembly comprises a chip on which an array of reactive species is immobilised, the chip being located in a storage well having a base and side walls. A protective, removable packaging may be provided over the storage well for protecting the assay device in transit.

This invention overcomes the problems mentioned above by placing the chip in a storage well, thus protecting the assay device while it is being handled during an immunoassay process. In addition, by providing a protective, removable packaging over the storage well (as is preferred), the complete assembly of chip, storage well, and packaging can be prepared centrally and then sent to the end user easily, without risk of damaging the chip and in particular the reactive sites. The form of the storage well will depend on a number of factors. If, for example, the reactive sites are to be examined using a chemiluminescent (or fluorescent or other lighting machine) technique, it is advantageous to make the walls of the storage well dark, preferably black, to reduce reflections/scattering of light and, where the storage well is joined to adjacent storage wells, to reduce or eliminate the transmission of light into adjacent wells. The base and sidewall of the storage well may therefore be made of a black material such as pigmented plastic or could be coated/painted in a black material.

The base is preferably continuous but could have a central aperture surrounded by a lip.

Another problem which can arise with the use of storage wells is due to the liquid meniscus of the liquid reagent which is used in the storage well during the immunoassay. If this reagent must remain in the well during a final analysis stage, the meniscus will contribute to distortion and aberration of the image of the reactive sites viewed by the camera system.

One approach is to place the chip on its edge in the storage well. The well would then need to have a transparent side wall to enable the reactive sites to be viewed or could have a sufficiently large top opening to enable the device to be viewed from the top.

Conventionally, however, the inner surface of the sidewall tapers inwardly adjacent the base. The use of a tapering sidewall allows the cross-sectional area of the open part of a well to be maximised and therefore that of the meniscus which is thereby flattened and thus the aberrations are reduced. Also, the chip can be laid flat on the base for viewing from the top.

Further flattening can be achieved by the selection of suitable well material and internal surface finish. Of course, the material chosen for the wells and any coatings applied to the inside should be chemically unreactive so as not to effect the immunoassay. Preferred materials comprise PVC and polypropylene.

A further advantage of the use of a taper is that it facilitates easy substrate placement and location. This is particularly important in the case of an automated process for loading substrates into the wells.

Although individual storage wells could be provided, preferably a number of such storage wells are provided fixed together in an array. This again simplifies the handling of chips by protecting them within the wells and also makes it easier to handle the storage wells since the array will have a larger size than each individual well.

Preferably, the chip is retained in the storage well by some form of retaining means. The retaining means could be in the form of retaining clips or adhesives to glue the substrates to the base. Neither of these is particularly desirable since they could effect the immunoassay. Preferably, the retaining means comprises one or more hot or cold formed projections on the inner surface of the sidewall. These could be formed prior to supplying the

substrate, which is then press fitted into the well, or after the substrate has been supplied.

Typically, each storage well is square in plan since this is suited to the square format of conventional CCD cameras. However, other plan forms such as rectangular or circular are envisaged.

To further ease handling, preferably the assembly further comprises a carrying tray for carrying one or more storage wells for use with an assay device processing instrument.

Such a carrying tray can then be used not only for holding the storage well(s) during supply to a user but also in an immunoassay machine.

15 BRIEF DESCRIPTION OF THE DRAWINGS

An example of an array of storage wells according to the invention will now be described with reference to the accompanying drawings, in which:-

Figure 1 is a perspective view of the array from above;

Figure 2 is a section taken on the line 2-2 in Figure 1 but showing a biochip in one of the storage wells;

Figure 3 is a perspective view of the section shown in Figure 2; and,

Figure 4 is a perspective view of a carrying tray for the array of storage wells.

DETAILED DESCRIPTION OF THE EMBODIMENT

Figure 1 illustrates an array of three storage wells 1-3 formed from a one-piece plastics moulding of P.V.C. or polypropylene. For the reasons given above, the plastics material incorporates a black pigment. Each storage well 1-3 has a similar form and as can be seen in Figure 1 is substantially square in plan. For convenience, only the storage well 1 will be described in detail.

The storage well 1 has a base 4 and a sidewall 5 surrounding the base. As can be seen in Figure 2, the

sidewalls 5 of each storage well are integrally formed at the junctions between the storage wells.

Protrusions 6 are moulded at each end of the array to enable the array to be handled easily.

5 Each sidewall 5 has an upper section 7 which is substantially vertical with respect to the base 4 and a lower section 8 which tapers inwardly. The taper terminates just short of the base 4 so as to define a region 9 having a width and height corresponding to that of
10 a biochip 10. Typical array dimensions are: 42mm long, 9mm high and 14mm wide at the top.

Following construction of the array of storage wells 1-3, each is supplied with a biochip 10. The biochips 10 can be prepared in any conventional manner so as to attach
15 ligands on respective reaction sites. For example, ligands could be immobilized by means of microfluidic dispensing of the ligand onto the substrate, which is chemically activated. Alternative chemical or physical methods could be used. It is important that the method of
20 immobilisation, e.g. covalent immobilisation, is such that ligands are not released during incubation and washing steps. Each chip which has dimensions 10mm x10mm and is about 1mm thick is then dropped into the respective storage well 1-3 and one such biochip 10 is shown in the storage
25 well 1 in Figures 2 and 3.

Each biochip 10 is then secured in the base of the storage well by cold or hot forming bumps 11 on at least one side section of the sidewall 5. These bumps may be either preformed for press fitting or post-formed after
30 insertion of the biochip 10.

As well as being tapered, the inner surfaces of the sidewalls 5 are preferably provided with a polished finish to reduce the curvature of the liquid meniscus and minimise optical aberrations.

35 Following these steps, the set of three storage wells can then be prepicked in an individual sealed "bubble" on a tape forming a roll for reel dispensing. However, in the

preferred approach, three sets of storage well arrays of the type shown in Figure 2 are loaded onto a carrying tray 20 as shown in Figure 4. This carrying tray is made of a plastics moulding and has two sets of crossbars 21,22 extending between opposite sidewalls 23,24 respectively. Nine openings 25 are defined into which the respective storage wells can be located. Each set of three storage wells 1-3 is loaded parallel to the crossbars 21 with the crossbars 22 entering into corresponding recesses 30 between adjacent storage wells. The loaded carrier tray is then sealed in suitable packing materials for transportation. The user can then either remove the storage wells from the carrier tray or, preferably, leave them in place and use the carrier tray to move the storage wells about the immunoassay process, for example as described in more detail in our copending European Patent Application No. 98307706.6.

A further option is to locate a number of the trays shown in Figure 4 with loaded storage wells into individual compartments of a stack defined by a housing. That housing can then be packaged for transportation. In this case, the trays could be directly extracted from the housing by an assay instrument or, of course, manually extracted as required.